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(54) TISSUE-SPECIFIC PROMOTER

(57) The invention provides a promoter capable of expressing an introduced gene in plant seeds, a vector comprising said promoter, a method for producing transgenic plants through transformation of plants with said vector, and a transgenic plant as transformed with said vector.

After a suitable foreign gene and a terminator are linked to the promoter, the resulting vector may be introduced into seeds of barley or other plants, thereby intentionally modifying the seeds and making the resulting seeds produce foreign substances therein.

Description

Field of Techonolgy

The present invention relates to a tissue-specific promoter, and particularly to a promoter for genes capable of being specifically expressed in plant seeds.

Background Technology

Barley which is one example of plants is an essential agricultural crop for feed and for producing food and drink (beer, whisky, etc.), and is worldwide cultivated and consumed. In accordance with many uses for barley, various breeding in barley have heretofore been made. One conventional breeding in barley comprises selecting some effective varieties from artificial or natural mutants followed by combining them through mating or the like to thereby find out from a number of the resulting progeny the hybrids capable of expressing the intended phenotype. However, as comprising mating, the breeding of this type is problematic in that the genotype to be introduced is limited to one for relative mating and that it takes a long period of time to obtain the intended hybrids.

On the other hand, with the recent development in biotechnology such as genetic engineering technology and cell technology, a system for directly introducing a desirable gene into plants is being established even for barley, and is expected to be one capable of overcoming the problems in the conventional breeding (for example, BIOTECHNOLOGY 13, 248, 1995 is referred to for barley for brewing). The system requires a tissue-specific promoter. In this, precisely, where a foreign gene is introduced into a plant, the gene is required to be sufficiently expressed in the intended tissue in good time. For this purpose, a tissue-specific promoter must be linked to the foreign gene to thereby make the gene expressible under the control of the promoter.

The present invention is to provide a promoter capable of being specifically expressed in plant seeds. As one example, we, the inventors of the present invention have succeeded in the isolation of a promoter region that acts to control the transcript ion of a barley β -amylase gene and also in the analysis of the nucleotide sequence of the promoter region.

Barley β -amylase is a β -amylase obtainable from barley seeds (1,4- α -D-glucanmaltohydrolase [EC 3.2.1.2]), which is known as an enzyme usable, like soybean β-amylase, in the industrial production of maltose for injection and maltose for food and drink. It is also known that barley may be germinated to give malt which may be a raw material for beer and liquors. β-amylase existing in malt is one of the most important enzymes for the saccharification of starch in the step of

Regarding the gene of barley β -amylase, the complete sequence of cDNA of a variety of barley, Hiproly, which comprises 1754 bases, has been reported, and the amino acid sequence thereof comprising 535 residues has also been deduced (see Eur. J. Biochem., 169, 517, 1987).

In addition, the complete sequence of cDNA of a variety of barley. Haruna Nijo, which comprises 1775 bases, has been reported, and the amino acid sequence thereof comprising 535 residues has also been deduced (see J. Biochem., 115, 47, 1994; Japanese Patent Application Laid-Open No. 6-303983). Further, the complete sequence of the structural gene region of the chromosome DNA of the variety, Haruna Nijo, which comprises 3825 bases, has been reported (see Japanese Patent Application No. 7-92004).

However, regarding the promoter region that acts to control the transcription of such a β -amylase gene, there is no report referring to the isolation of the promoter gene, let alone the analysis of the nucleotide sequence thereof.

We, the present inventors, having attempted to isolate the promoter region of a barley β-amylase gene that may be actively expressed in barley developing seeds, thereby utilizing it in the improvement of barley seeds or in the production of products in barley seeds, have earnestly studied to attain this object, and, as a result, have completed the

According to the present invention, a desired foreign gene and a terminator therefor can be linked to the downpresent invention. stream site of the promoter region obtained, and introduced into plant such as barley, whereby the foreign gene can be expressed in the plant developing seeds. Thus, the promoter region can be utilized in the improvement of seeds of barley and other plants and also in the production of substances in such seeds.

Disclosure of the Invention

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The first aspect of the present invention is a promoter that acts to express an introduced gene in plant seeds.

The promoter includes, for example, one which has a molecular weight of 1.28 kb and which comprises cleavage sites to be cleaved with restriction enzymes, Sal I, Apa I, Xba I, Bam HI, Hind III, EcoT 221, Xba I and Bam HI, in that order (this corresponds to the white area shown in Fig. 2), and one which comprises a gene substantially having the nucleotide sequence of Sequence Number 1 in Sequence Listing.

The second aspect of the present invention is a vector comprising said promoter.

The third aspect of the present invention is a method for producing transgenic plants, which comprises transforma-

tion of plants with said vector.

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The fourth aspect of the present invention is a transgenic plant as transformed with said vector.

Brief Description of the Drawings

Fig. 1 is a physical map showing a clone that comprises the promoter region of a β-amylase gene.

Fig. 2 is a physical map showing a 2.4 kb Sal I-Sal I fragment that comprises the 5'-terminal region of a structural gene of a β -amylase gene and the promoter region of the β -amylase gene.

Fig. 3 shows a process for constructing a reporter plasmid.

Fig. 4 is a graph showing the GUS activity of various cell lines.

Fig. 5 shows an expression vector pSBG503 of a thermophilic β-amylase.

Fig. 6 is a graph comparatively showing the thermophilicity of various β -amylases as extracted from the seeds of plant individuals derived from protoplasts into which was introduced a thermophilic β -amylase gene according to the present invention.

Best Mode to practice the Invention

As has been mentioned hereinabove, barley β -amylase is a β -amylase (1,4- α -D-glucanmaltohydrolase [EC 3.2.1.2]) to be obtained from barley seeds.

Of the gene of this enzyme, the complete sequence of cDNA as derived from particular varieties of barley has been reported, as has been mentioned hereinabove. However, regarding the promoter region that acts to control the transcript ion of such a β -amylase gene, there is as yet no report referring to the isolation of the promoter gene and to the analysis of the nucleotide sequence thereof.

It is known that, in barley plants, barley β-amylase is specifically produced in the developing seeds, and that the enzyme is an essential protein that accounts for approximately from 1 to 2 % of the soluble proteins in the endosperm (see Hereditas, 93, 311, 1980).

Having known the above, we, the present inventors expected that a promoter region capable of being specifically expressed in plant seeds could be utilized as the transcriptional control factor for a foreign gene to be introduced into plant for the purpose of making the plant seeds produce the foreign gene-related substance. In addition, for barley, we specifically noted that the isolation as well as the identification of the promoter region of a barley β-amylase gene is important in order to successfully introduce a foreign gene into barley by the use of the promoter.

To concretely illustrate the present invention, one process is described in detail hereinunder which comprises isolating and analyzing the promoter region of a barley β -amylase gene, constructing an expression vector that comprises the promoter region, introducing the expression vector into plant along with a foreign gene, and making the thus-transformed seeds express the foreign gene.

The "gene substantially having the nucleotide sequence of Sequence Number 1 in Sequence Listing" as referred to herein means that the gene allows any deletion, substitution and addition of some bases in said sequence so far as the gene has the significant promoter activity in plant seeds.

(1) Preparation of Barley Chromosome DNA:

In barley, the same barley chromosome DNA exists in all cells of every tissue. From barley seeds as germinated in vermiculite in the dark at 20°C for 7 days, the primary leaves can be processed herein to give the intended barley chromosome DNA. The preparation of the DNA can be conducted by any known method. For example, referred to is the method described in "Cloning and Sequencing - Manuals for Experiments in Plant Biotechnology" (published by Nohson Bunka Publishing Co., 1989), page 252.

(2) Formation of Barley Genomic Library:

Using the barley chromosome DNA, a barley genomic library can be formed by any known method. For example, referred to is the method described in "Cloning and Sequencing - Manuals for Experiments in Plant Biotechnology" (published by Nohson Bunka Publishing Co., 1989), page 272.

(3) Formation of Probe:

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Probes to be used for screening the barley genomic library can be formed by labeling a suitable DNA fragment, or that is, a DNA fragment having a sequence that is complementary to the sequence of the gene to be selected through the screening, with DIG-High Prime (produced by Boehringer Mannheim Co.).

(4) Cloning of Promoter Region of Barley β-amylase Gene:

To clone the promoter region of the barley β-amylase, the barley genomic library may be screened by the use of the probe as formed in the above (3). This screening can be effected by any known method. For example, referred to is the method described in "Cloning and Sequencing - Manuals for Experiments in Plant Biotechnology" (published by Nohson Bunka Publishing Co., 1989), page 134. The detection of the intended clones can be effected by the use of DIG Luminescent Detection Kit (produced by Boehringer Mannheim Co.).

(5) Sequencing:

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The promoter region can be sequenced, for example, according to the Maxam-Gilbert chemical degradation of DNA method (see Methods in Enzymology, 65, 499, 1980) or the Sanger dideoxy-mediated chain-termination method (see Gene, 19, 269, 1982).

5 (6) Constrution of Reporter Plasmid:

To determine the promoter activity of the promoter region thus obtained through the steps mentioned above, a reporter gene, such as a β -glucionidase gene (GUS), and a terminator, such as a nopalin synthetase gene (NOS) terminator, may be linked to the downstream site of the promoter region to form a reporter plasmid, and the activity of the product translated from the reporter gene may be measured. As the reporter gene and the terminator, usable are commercially-available products, such as plasmid pBI 101 (produced by Clontech Co.).

(7) Detection of Promoter Activity in Endosperm Cells of Developing Seeds:

To determine the promoter activity in endosperm cells of developing seeds by the use of the reporter plasmid as constructed in the above, any known method can be employed (see Plant Cell Reports, 10, 595, 1992). Briefly, a protoplast is prepared from endosperm cells of developing seeds, into which is introduced the reporter plasmid according to a known method, for example, a polyethylene glycol method (see, for example, Theor. Appl. Genet., 91, 707, 1995; Japanese Patent Application Laid-Open No. 7-184492), and the GUS activity in the resulting cell lines is measured.

(8) Formation of Transgenic Plant:

Using the promoter of a barley β -amylase gene of the present invention, an expression vector is constructed. Then, the vector is introduced into plant cells to obtain a transgenic plant.

As one example, the expression vector plasmid for use in the present invention comprises a thermophilic β -amylase gene as the gene to be expressed, the promoter of the invention as the transcriptional control factor, and a cauliflower mosaic virus 35S terminator as the terminator. This expression vector plasmid is introduced into plant, which thereby can produce the intended thermophilic β -amylase in seeds.

The thermophilic β -amylase gene to be linked to the promoter may be any organism-derived one or may even be any modified one to be prepared by modifying the organism-derived gene. We, the present inventors employed herein a thermophilic β -amylase gene as obtained through site-specific mutation of a barley β -amylase gene (see Japanese Patent Application Laid-Open No. 7-327681).

In order to directly introduce the recombinant plasmid into plant cells, employable is any of electroporation methods (for example, see Nature, 319, 791, 1986), polyethylene glycol methods, particle gun methods (for example, see Nature, 327, 70, 1987), laser perforation methods (for example, see Barley Genetics VI, 231, 1991), Agrobacterium methods (for example, see Plant J., 6, 271, 1994) and others. We, the present inventors employed herein barley as the test material and a polyethylene glycol method using protoplasts as the gene introduction method.

Barley protoplasts can be prepared preferably from an immature embryo-derived callus (see Kihara & Funatsuki, 1995, Plant Sci., 106: 115-120; Japanese Patent Application Laid-Open No. 7-213183) or from suspension culture cells with regeneration ability as established from such an immature embryo-derived callus (see Kihara & Funatsuki, 1994, Breeding Sci., 44: 157-160; Funatsuki & Kihara, 1994, Plant Cell Rep., 13: 551-555; Japanese Patent Application Laid-Open No. 4-360633), according to any ordinary protoplast preparation method using cellulase and pectinase.

After the formation of colonies from the protoplasts, the liquid medium and the cell suspension used as nurse cells are removed, and the colonies are further cultured in a liquid medium containing a selective reagent, such as geneticin (G418), hygromycin, bialafos or the like. Thus, only resistant colonies grow in the medium.

The thus-grown colonies are transferred onto a solid medium containing any of the selective reagents, geneticin (G418), hygromycin, bialafos and others. Further culture on the solid medium gives embryogenic calluses or embryoids, which are then transferred onto a different solid medium containing no selective reagent, resulting in their regeneration into plant.

The thus-grown plant individuals are then transplanted in pots and are cultivated therein under ordinary cultivation conditions, for example, at a daylength of 16 hours, at 10,000 luxes and at 18°C, thereby being fertile transgenic plants.

From the seeds obtained from the plants, extracted is a β -amylase, which is then heat-treated. The activity of the thus heat-treated enzyme is measured to determine its thermophilicity. The amylase activity can be measured through saccharification of starch with the enzyme. We, the present inventors employed herein an amylase determination reagent (Diacolor AMY, trade name of a product of Ono Pharmaceutical Co.), with which only the activity of β -amylase can be selectively determined even in a small amount of a sample containing the enzyme.

Examples

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Now, the present invention will be described in detail hereinunder with reference to the following examples, which, however, are not intended to restrict the scope of the present invention.

Example 1:

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Preparation of Barley Chromosome DNA:

About 1000 grains of barley (Haruna Nijo) were germinated in vermiculite in the dark at 20°C for 7 days. The primary leaves (about 65 g) thus grown were taken off and then cut into fine pieces of about 1 cm long, from which was prepared a chromosome DNA. As a result, about 1 mg of DNA was extracted from 10 g of the leaves.

Example 2:

Formation of Barley Genomic Library:

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150 μg of the chromosome DNA as prepared in Example 1 was partially digested with 1 U of Sau 3AI, at 37°C for 1 hour, and the resulting fragments were fractionated according to sucrose density gradient centrifugation. The fraction comprising fragments of about 18 kb was purified and inserted into a λ phage vector EMBL3 (produced by Stratagene Co.). Using Gigapack II Gold (produced by Stratagene Co.), the resulting vector was packaged into lambda phage particles, with which Escherichia coli XL1-Blue MRA(P2) (produced by Stratagene Co.) were transformed.

Example 3:

Formation of Probe:

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A barley β-amylase structural gene-derived Eco RV-Hind III fragment described in Japanese Patent Application No. 7-92004, or that is, the DNA fragment having the nucleotide sequence of Sequence Number 2 in Sequence Listing was labeled with digoxigenin, using DIG-High Prime (produced by Boehringer Mannheim Co.), to obtain a probe.

40 Example 4:

Cloning of Promoter Region of Barley β -amylase Gene:

The plaque of the barley genomic library as formed in Example 2 was transferred onto a nylon membrane, "Hybond N" (produced by Amersham Co.), and then screened through ordinary plaque hybridization using the probe as formed in Example 3. To detect the intended clone, used was DIG Luminescent Detection Kit (produced by Boehringer Mannheim Co.). As a result, one positive clone was obtained. This clone had the 5' - terminal region of the β-amylase structural gene and the upstream region containing a promoter region for the gene.

The physical map of the thus-obtained clone is shown in Fig. 1, in which the abbreviations indicate the sites that are recognized and cleaved by the indicated restriction enzyme, the thin lines indicates the vector sites, the black area indicates the 5'-terminal region of the β -amylase structural gene, and the white area indicates the upstream site containing the promoter region. The arrow therein indicates the direction of the β -amylase gene.

Example 5:

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Sequencing of Promoter Region of β -amylase Gene:

From the positive clone as obtained in Example 4, cleaved out was the Sal I-Sal I fragment of 2.4 kb composed of the 5'-terminal region of the β -amylase structural gene and the promoter region. This fragment was inserted into a plas-

mid pUC119, from which was formed a deletion clone using Kilo-Sequence Deletion Kit (produced by Takara Shuzo Co.). After this, the promoter region was sequenced according to Sanger dideoxy-mediated chain-termination method.

The physical map of the 2.4 kb Sal I-Sal I fragment is shown in Fig. 2, in which the abbreviations indicate the sites that are recognized and cleaved by the indicated restriction enzymes, the black area indicates the 5'-terminal region of the β -amylase structural gene, and the white area indicates the upstream site containing the promoter region.

The nucleotide sequence of the thus-sequenced promoter region of the β -amylase gene is Sequence Number 1 in Sequence Listing. The partial nucleotide sequence of the 2.4 kb Sal I-Sal I fragment thus sequenced is Sequence Number 3 in Sequence Listing. Comparing the nucleotide sequence of the 5'-terminal region of the β -amylase structural gene with the barley β -amylase structural gene that had already been obtained (see Japanese Patent Application No. 7-92004), it was confirmed that the DNA fragment obtained herein is a β -amylase gene. The promoter region sequenced herein contained a TATA box which widely exists in promoter regions in eucaryotes.

Example 6:

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5 Formation of Reporter Plasmid:

A reporter plasmid was formed in accordance with the method illustrated in Fig. 3. Precisely, a Hind III-Eco RI fragment comprising the GUS gene and the NOS terminator of a plasmid pBI 101 (produced by Clontech Co.) was inserted into the Hind III-Eco RI site of a plasmid pUC 118 to prepare a plasmid pBI 11.

On the other hand, of the fragment of Sequence Number 3 which is comprised of from the 1st to the 1672nd bases of the nucleotide sequence of the deletion clone as formed in Example 5, or that is, the plasmid composed of the promoter region of Sequence Number 1 and the 341 bp 5' - terminal region of the β -amylase structural gene, cleaved out was a Pst I-Eco RI fragment containing said promoter region. The ends of the thus-cleaved Pst I-Eco RI fragment were blunted, using a blunting kit (produced by Takara Shuzo Co.), and the thus-blunted fragment was inserted into the Sma I site of the plasmid pBI 11 to give a reporter plasmid pSBG 530.

Further, the Hind III-Hind III fragment that codes for the 5' - terminal side of the β-amylase promoter region of the plasmid pSBG 530 was removed from the plasmid to give another reporter plasmid pSBG 530dH.

Example 7:

Detection of Promoter Activity in Endosperm Cells in Developing Seeds:

The activity of the isolated promoter region of the β -amylase gene in endosperm cells in developing seeds was determined in a transient assay system using the reporter plasmids formed in Example 6.

First, developing seeds of a variety of barley, Bomi that had been harvested in about 14 days from the blooming were peeled to remove their husks, then sterilized once with 70 % ethanol and once again with a 1/5 dilution of hypochlorous acid, and thereafter washed water for a total of three times. The endosperm was extracted out from these, and processed overnight with a CPW solution (0.2 mM KH₂PO₄, 10 mM CaCl₂, 1 mM MgSO₄, 1 mM KNO₃) containing 0.4 % cellulase and 11% mannitol, at 25 °C.

The resulting protoplasts were washed with the CPW solution containing 11% mannitol, and then divided into plural sections of 10^6 protoplasts each per one transformation system. To each protoplast section, added were 30 μ g of the DNA and 200 μ l of a C100S solution (7% sorbitol, 10 mM CaCl₂, 4.7 mM MES, pH 5.7) and suspended. The resulting suspension was then processed with 0.5 ml of the C100S solution (pH 7.0) containing 40 % polyethylene glycol 1540 added thereto, for 10 minutes.

To this was added 10 ml of an LW solution (see Lazzeri et al., Theor, Appl, Genet., 81:437, 1991), and the resulting mixture was centrifuged. 3 ml of an L1 medium (see Theor, Appl, Genet., 81:437, 1991) was added to the resulting residue, which was then incubated overnight at 25°C. 20 ml of the LW solution was added to the resulting culture, which was then centrifuged. The resulting residue was suspended in 200 µl of a GUS extract (0.05 M NaPO₄, 0.01 M EDTA, 0.1% sarcosine, 0.1 % Triton X-100. 0.1 % 2-mercaptoethanol), and the suspension was then frozen and thawed repeatedly twice. This was centrifuged, and the resulting supernatant was used as a crude enzyme solution for the determination of the promoter activity.

Precisely, the crude enzyme solution obtained hereinabove was reacted with 4-methylumbelliferyl-β-D-gluclonide, then the reaction was stopped with 0.2 M sodium carbonate solution, and the 4-methylumbelliferyl residue produced was quantified, from which was determined the promoter activity. To quantify the protein, used was "Protein Assay" produced by Bio-Rad Co.

The GUS activity in each cell line is shown in Fig. 4, from which it is confirmed that the isolated β-amylase promoter region was active in the endosperm cells of developing barley seeds. The GUS activity in the cell line introduced pSBG 530dH was lowered to about 2/3 of that in the cell line introduced pSBG 530, from which it is confirmed that the promoter region requires the nucleotide sequence of Sequence Number 1 in Sequence Listing.

Example 8:

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Formation of Transgenic Plant:

In accordance with the method of Kihara & Funatsuki (1994, Breeding Sci., 44:157-160) or the method of Funatsuki & Kihara (1994, Plant Cell Rep., 13:551-555), immature embryos having a length of approximately from 0.5 to 1.0 mm of a variety of barley, Igri, were placed onto L2 medium for callus indication. After one month, the thus-formed calluses were transferred onto an L1 liquid medium and cultured therein for from 2 to 4 months by shaking culture, while being exposed to weak light (at 500 luxes). Thus was formed a liquid suspension culture comprising cell masses having a diameter of approximately from 1 to 3 mm.

To 1 g of the cells, added were about 10 ml of an enzyme solution (1.0 % Cellulase Onozuka RS, 0.1 % Pectolyase Y-23, 5 mM MES dissolved in LW solution) and the resulting mixture was left statically at 25°C for 2 to 3 hours.

The thus-obtained protoplast suspension was filtered through a 64- μ mesh membrane and a 26- μ mesh membrane, and then centrifuged to collect the protoplasts. Then, these were washed with an LW solution for a total of three times

Next, from 1 x 10^6 to 3 x 10^6 protoplasts thus obtained were suspended in 250 μ l of a liquid medium, Ca-S which comprised 10 μ g/ml of a plasmid pSBG503 (expression vector for thermophilic β -amylase - see Fig. 5), 100 mM of CaCl₂, 0.6 M of sorbitol and 0.1 % of MES and which had been adjusted to pH 5.7. The plasmid pSBG503 comprised a kanamycin-resistant gene and a thermophilic β -amylase gene, in which a rice actin promoter and a cauliflower mosaic virus 35S terminator (35 St) were linked to the kanamycin-resistant gene while the barley β -amylase promoter region of Sequence Number 1 and a cauliflower mosaic virus 35S terminator were to the thermophilic β -amylase gene, each as the transcriptional control factor and the terminator, respectively. The thermophilic β -amylase gene comprised the first intron of a barley β -amylase gene. To the resulting suspension was dropwise added 600 μ l of Ca-S which contained 40 % of polyethylene glycol and which had been adjusted to pH 7.0. This was statically left as it was for 10 minutes, while being shaken at intervals of 5 minutes. This was diluted with 10 ml of an LW solution and then centrifuged to collect the protoplasts.

The thus-collected protoplasts were then suspended in 1 ml L1 medium containing 0.6 M maltose, 2.0 mg/liter 2,4-D and 1.8% agarose, and immediately spread over a 6-cm Petri dish to make thereon a disc having a diameter of about 4.5 cm. After having been solidified, the resulting solid was peeled off from the dish, and then incubated in 5 ml of a liquid medium (this comprised of the same components as those constituting the medium used hereinabove to make the protoplast suspension) which contained 200 mg/ml of barley suspension cells, with shaking at a shaking speed of 50 rpm.

On 15 days after the start of the culture of the protoplasts, the liquid medium and the suspension cells were removed, and 3 ml of a liquid medium containing 20 µg/ml of Geneticin (G418) was added to the protoplast culture. Then, the resulting protoplast culture was further cultured for 14 days with shaking, resulting in the growth of resistant colonies in agarose and therearound and also in the liquid medium.

The thus-grown colonies were transferred onto L3 medium containing 20 µg/ml Geneticin (G418) and containing, as hormones, 0.5 mg/liter of 2,4-D and 1.0 mg/liter benzylaminopurine (BAP). On 3 to 15 days after the transfer, embryogenic calluses or embryoids were found to grow on the selection medium.

These calluses or embryoids were transferred onto L3 medium not containing any selective reagent but containing 0.5 mg/liter 2,4-D and 1.0 mg/liter BAP. To this stage, the incubation was conducted under weak lighting (at about 500 luxes) at 25°C. After about 3 to 15 days, regeneration of shoots was observed from the calluses or embryoids. After having been sufficiently shoot development, these were transferred into a light place where they were exposed to strong light of about 7000 luxes.

The thus-grown barley plantlets were then transplanted onto L3 medium not containing any hormone, for the induction of root development. After about one month, these were transplanted in pots. After having been thus transplanted in pots, these were cultivated therein at a daylength of 16 hours, at 10,000 luxes and at 15°C, thereby being a large number of transgenic barley plants. The presence or absence of the thermophilic β-amylase gene fragment in the thusgrown barley plants was checked through polymerase chain reaction (PCR) which verified the presence of the fragment therein.

From the developing seeds these barley plants, extracted was an amylase using 50 mM acetate buffer containing 10 mM DTT. The thus-extracted enzyme was heat-treated at temperatures falling between 50 and 75°C (varying at intervals of 2.5 °C) for 30 minutes, and the β-amylase activity of the enzyme was measured, using Diacolor AMY, to determine its thermophilicity. The results are shown in Fig. 6.

As is obvious from the data in Fig. 6, it was verified that the seed samples, a and b, both having the thermophilic β -amylase gene as introduced thereinto contained the intended thermophilic β -amylase as accumulated therein, while the control seed sample, p, derived from the protoplasts not having the thermophilic β -amylase gene did not contain it. In particular, it is known that the accumulation of the thermophilic β -amylase in the seed sample, a, is remarkable.

Possibility of Industrial Utilization

According to the present invention, there is provided a promoter for a gene capable of being specifically expressed in plant seeds. In particular, the present invention has clarified the nucleotide sequence of the promoter region for a β -amylase gene and has clarified the activity of the promoter in developing seeds. After a suitable foreign gene and a terminator are linked to the promoter, the resulting vector may be introduced into seeds of barley or other plants, thereby intentionally modifying the seeds of barley or other plants. In addition, it is also possible to make the resulting transgenic plant seeds produce foreign substances therein. Thus, the present invention produces many advantages in the field of plant breeding.

SEQUENCE LISTING

,	Sequence Number: 1								
	Length of sequence: 1276								
10	Type: nucleic acid								
	Strandedness: double								
	Topology: linear								
	Molecule Type: DNA								
15	Sequence:								
	GTCGACACAT CATCTTGAGA ACGTCTTCTC TAACTGAATG CCAATTGCTG AAAGGCATGA	60							
20	AATAAATTAA GGAAACTTGC AAACACAAAT GTCAAGTCTG AGACTTGAAC TCTAGTGGAC	120							
	ACGTTTGAAC ATTAATGTGC GGTTTTGGTG AAAGTGAAAA AACAGTTGAC ATCCTTGTGA	180							
	ATGAATCAAT TCAAACCAAA TGCAAAAATG AGAAGTGAAA ACCCCATAAT CGTCGTTTGT	240							
25	AGAAAACAAA TAACACTGAA ACTAATGGGG TAATTTCTGA GAAACTATAA TTGGTGAGGC	300							
	ACATTCTCAT TTGATTGTTT AGTTTAACTT CCTTCTCACA TTATTTTTTC CTAGAGTATC	360							
30	TCATTGCTCG GGTGGCATCC AAATTTTCCA GACAAGGGCC CATGCAATGG TTCTAGATGA	420							
	TTGGAAAACA AATATTAAAT ATTTTTGTAG ATGATTGTAT TAGTGTCGCA AACATGCAAC	480							
	ACAATTTTCG TCGGAAAAAA AAACAAATTT AGGATGACAT TTTGGGGTAA CTTTTGGTGT	540							
35	TCAATTTGTT TTTTTGGCAC AAGCCAAAAT GTTTAACCTT TTTACCTCAA AATTTGCAGG	600							
	TAGCATTTAG ATATGACTAA GTATATAAAT AAATTTTGTC TTGATTTTTT TTCATTTTGA	660							
	ATTTTTTGG CCCCCGAAGC ATATTCTTCC GGGAGCCAAA TTGACATTCC GGTCATGATG	720							
40	TGGCTTGGAT CCCAAGTTAG TCATACAGAT AAGGATATAT CTTACCTCAA CCGAATCTAG	780							
	GTTACAACAA GCTTAACACT CATGCATTAG TGTCCATCCT AGACTCCTCT AGAAGGCAAT	840							
45	GGTTTACACA CCATTGTGGT TTGTACAAAT TCCAACACTC TTCCTCAATC ATAACTTTAT	900							
	GAAGTCGAGA TTGATCTTAA AGTTCTGATG TTATCTGTTA AAGAATAGCT TGGGAAACAC	960							
50	ATGTACAACT TAAGTCAGGA TGTGTGCGTC TTCACTTCGT ATAGGGTGCC GTTTGGTTGA	1020							
	GAGTTGAGAC GTGGATTGGA TCCCACGTTA GCTATAATAT AGATAATGAT CCCATCTCTC	1080							
	TCCCAATCAG ACCTCAATCC TTGAAGTTTG GCTGTGTGTG GTAAAGAAAA CCAAAATTAG	1140							

	TTCACAAAGC GO	CCCTTTCTG	AGTGGACAAT	CCTATTTCTG	ACATATCTGA	TTGGAAAGCT*	1200		
5	AGGTTCGCCG TT	TGGCCTCAC	ATCTATGGAT	ACATCTTTTT	CATAGTATAA	ATAGACCCTT	1260		
	TTATTAAGCT CO	CCTGC					1276		
10									
	Sequence Numb	ber: 2							
	Length of sequence: 1066								
15	Type: nucleic acid								
	Strandedness	: double							
20	Topology: linear								
	Molecule Type	e: DNA					•		
	Sequence:								
25	GATATCCAAC A						60		
	GGTGAACGTG A						120		
30	TTCAGACCAA T						180		
	TACATTATTG T						240		
	AGAACAAGAT C						300		
35	ATTGTTGAAA T						360		
	GACGAGCTGA G						420		
40	GACGTCTGGT G						480		
	AAGCAGTTGT T						540		
	CACCAGTGTG G						600		
45	GACGTCGGCA C						660		
	GAGTACCTCA C						720		
50	GTTACTTTAA A						780		
50	ACCAAATACA A	GCAAAAGGT	TAAAGGTGCC	AAAAACAGAT	AAGCAAAGAA	ACAAAACCTA	840		
	CCTAATCAAA C	CACTCTAGAG	CCTATCAAAA	AAAAAAAAA	AACATCGAGA	AGGTGCCTAG	900		

	AGCGGATGGG '	T TTCGACAAC	CCTTTAGCTT	TCATGCATCT	TTTTGGGAAA	GGGTGAAAA'A `	960		
	CACCGTCCTT '	TAAGTCGATT	GATGCAGGCA	GCCTTCTATT	GTTTGTAAGC	TATCAGGAAA	1020		
5	TACAAAATTA	ATAGCTAGTT	GTCATTTTAA	TAGTTGTAGC	AAGCTT		1066		
10									
	Sequence Nu	mber: 3							
	Length of sequence: 2142								
15	Type: nucleic acid								
	Strandedness: double								
20	Topology: linear								
	Molecule Type: DNA								
	Sequence:					٠			
25	GTCGACACAT						60		
					AGACTTGAAC		120		
30						ATCCTTGTGA	180		
30 0		•				CGTCGTTTGT	240		
		•				TTGGTGAGGC	300		
35						CTAGAGTATC	360		
	TCATTGCTCG	GGTGGCATCC	AAATTTTCCA	GACAAGGGCC	CATGCAATGG	TTCTAGATGA	420		
	TTGGAAAACA	AATATTAAAT	ATTTTTGTAG	ATGATTGTAT	TAGTGTCGCA	AACATGCAAC	480		
40						CTTTTGGTGT	540		
	TCAATTTGTT	TTTTTGGCAC	AAGCCAAAAT	GTTTAACCTT	TTTACCTCAA	AATTTGCAGG	600		
45	TAGCATTTAG	ATATGACTAA	GTATATAAAT	AAATTTTGTC	TTGATTTTTT	TTCATTTTGA	660		
	ATTTTTTTGG	CCCCCGAAGC	ATATTCTTCC	GGGAGCCAAA	TTGACATTCC	GGTCATGATG	720		
	TGGCTTGGAT	CCCAAGTTAG	TCATACAGAT	AAGGATATAT	CTTACCTCAA	CCGAATCTAG	780		
50	GTTACAACAA	GCTTAACACT	CATGCATTAG	TGTCCATCCT	AGACTCCTCT	AGAAGGCAAT	840		
	GGTTTACACA	CCATTGTGGT	TTGTACAAAT	TCCAACACTO	TTCCTCAATO	ATAACTTTAT	900		

	GAAGTCGAGA	TTGATCTTAA	AGTTCTGATG	TTATCTGTTA	AAGAATAGCT	TGGGAAACAC	960
5	ATGTACAACT	TAAGTCAGGA	TGTGTGCGTC	TTCACTTCGT	ATAGGGTGCC	GTTTGGTTGA	1020
	GAGTTGAGAC	GTGGATTGGA	TCCCACGTTA	GCTATAATAT	AGATAATGAT	CCCATCTCTC	1080
	TCCCAATCAG	ACCTCAATCC	TTGAAGTTTG	GCTGTGTGTG	GTAAAGAAAA	CCAAAATTAG	1140
10	TTCACAAAGC	GCCCTTTCTG	AGTGGACAAT	CCTATTTCTG	ACATATCTGA	TTGGAAAGCT	1200
	AGGTTCGCCG	TTGGCCTCAC	ATCTATGGAT	ACATCTTTTT	CATAGTATAA	ATAGACCETT	1260
15	TTATTAAGCT	CCCTGCCATA	TCCAACAAAC	CATTTGAAGT	TGTAGAGCAT	CATCCATAGC	1320
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	CTGTAAGCTC	CATCCATTCA	GACCAATCGC	TGAGAACCAC	ACACTAAAAC	TATTTCAAGG	1440
20	ATCTAGTGCA	CACATATACA	TTATTGTTGT	ACATATAACA	TTGATACTTC	TTGTAAAACT	1500
	CTAATTCAAA	GGGTGAAGAA	CAAGATCTGA	GGCCTCAAAT	GAGTATTTTA	TTTGTACTAA	1560
25	CCTTGACTAC	ACTTCCATTG	TTGAAATAAA	TAAATAGCTG	GACGCCGTGA	GCGTGAACAA	1620
-	CAGGTTCGAG	AAGGGCGACG	AGCTGAGGGC	GCAATTGAGG	AAGCTGGTAG	AGGCCGGTGT	1680
30	GGATGGTGTC	ATGGTAGACG	TCTGGTGGG	CTTGGTGGAG	GGCAAGGGCC	CCAAGGCGTA	1740
	TGACTGGTCC	GCCTACAAGC	AGTTGTTTGA	GCTGGTGCAG	AAGGCTGGGC	TGAAGCTACA	1800
	GGCCATCATG	TCGTTCCACC	AGTGTGGTGG	CAACGTCGGC	GACGCCGTCA	ACATCCCAAT	1860
<i>35</i>	CCCACAGTGG	GTGCGGGACG	TCGGCACGCG	TGATCCCGAC	ATTTTCTACA	CCGACGGTCA	1920
	CGGGACTAGG	AACATTGAGT	ACCTCACTCT	TGGAGTTGAT	AACCAGCCTC	TCTTCCATGG	1980
40	AAGATCTGCC	GTCCAGGTTA	CTTTAAACCA	CCACTCTAGT	TCTCTGATGC	ATATTTATAT	2040
	AGAAGTTCAA	GATGACACCA	AATACAAGCA	AAAGGTTAAA	GGTGCCAAAA	ACAGATAAGC	2100
	AAAGAAACAA	AACCTAGCTA	ATGAAACAGT	CTAGAGCCTA	TC		2142

Claims

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- 1. A promoter capable of expressing an introduced gene in plant seeds.
- 2. The promoter as claimed in claim 1, wherein the expression of the introduced gene is possible in developing plant seeds.
- 3. The promoter as daimed in claim 1 or 2, wherein the expression of the introduced gene is possible in barley seeds.
- 4. The promoter as claimed in any one of claims 1 to 3, which has a molecular weight of 1.28 kb and comprises cleavage sites to be cleaved with restriction enzymes, Sal I, Apa I, Xba I, Bam HI, Hind III, EcoT 221, Xba I and Bam HI,

in that order (corresponding to the white area shown in Fig. 2).

- 5. The promoter as claimed in any one of claims 1 to 4, which comprises a gene substantially having the nucleotide sequence of Sequence Number 1 in Sequence Listing.
- 6. A vector comprising a promoter of any one of claims 1 to 5.

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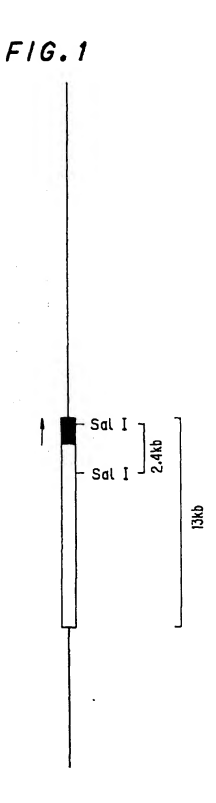
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- 7. A method for producing transgenic plants, which comprises transformation of plants with a vector of claim 6.
- 10 8. The method for producing transgenic plants as claimed in claim 7, wherein the introduced gene is a β-amylase gene.
 - 9. The method for producing transgenic plants as claimed in claim 8, wherein the β-amylase gene is a thermophilic β-amylase gene.
 - 10. The method for producing transgenic plants as claimed in any one of claims 7 to 9, wherein the transgenic plant is barley.
 - 11. A transgenic plant as transformed with a vector of claim 6.
 - 12. The transgenic plant as claimed in claim 11, wherein the introduced gene is a β -amylase gene.
 - 13. The transgenic plant as claimed in claim 12, wherein the β -amylase gene is a thermophilic β -amylase gene.
- 25 14. The transgenic plant as claimed in any one of claims 11 to 13, which is barley.



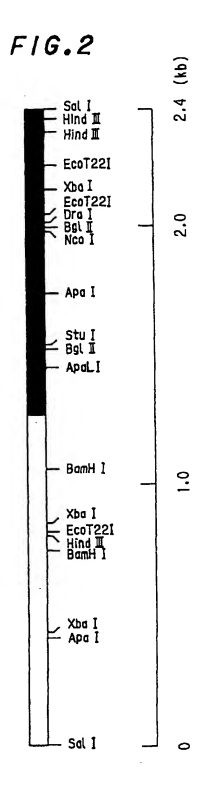


FIG.3

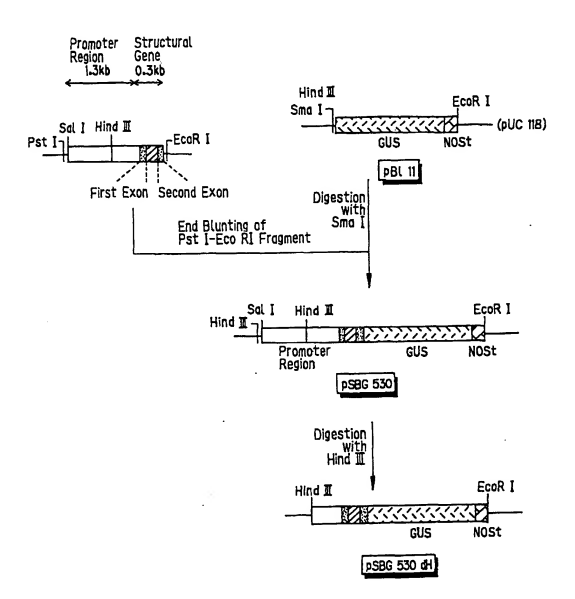
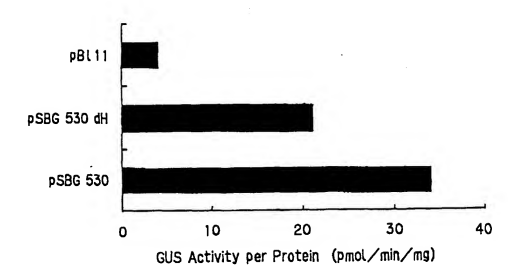
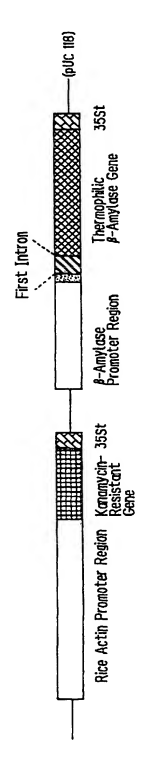


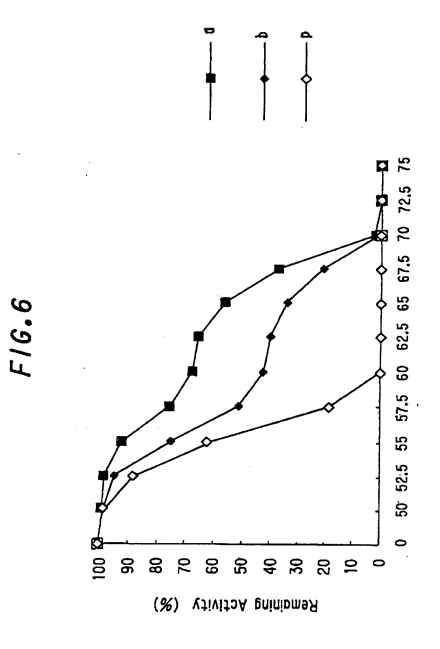
FIG.4



F16.5



pSBG 503



Heat-Treating Temperature (c)

19

INTERNATIONAL SEARCH REPORT International application No. PCT/JP96/01866 CLASSIFICATION OF SUBJECT MATTER Int. C1⁶ C12N15/82, A01H1/00, A01H5/00 // C12N5/14, (C12N15/82, C12R1:91), (C12N5/14, C12R1:91) According to International Patent Classification (IPC) or to both national classification and IPC FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) Int. C16 C12N15/82, A01H1/00, A01H5/00, C12N5/14 Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) WPI/L, BIOSIS PREVIEWS, CAS ONLINE C. DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriate, of the relevant passages Category* Relevant to claim No. X/Y JP, 7-143895, A (National Science Counsil), June 6, 1995 (06. 06. 95) 1-3, 6-7, 10-11, 14/ & US, 5460952, A 1-3, 6-14X/Y Proc. Natl. Acad. Sci. USA, Vol. 83, (1986), 1-3, 6 / 1-3, 6-14 R.N. Beachy et al., see p. 8560-8564 JP, 7-327681, A (Sapporo Breweries Ltd.), Y 1-3, 6-14 December 19, 1995 (19. 12. 95) (Family: none) J. Biochem., Vol. 115, (1994), S. Koshino et al., А 4 - 5 see p. 47-51 Further documents are listed in the continuation of Box C. See patent family annex. later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the inventors Special categories of cited documents: "A" document defining the general state of the art which is not considered to be of particular relevance "X" document of particular relevance; the claimed inveation cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone "E" earlier document but published on or after the international filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another clintion or other special reason (as specified) document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art "O" document referring to an oral disclosure, use, exhibition or other document published prior to the international filling date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report September 24, 1996 (24. 09. 96) October 15, 1996 (15. 10. 96) Name and mailing address of the ISA/ Authorized officer Japanese Patent Office Facsimile No. Telephone No. Form PCT/ISA/210 (second sheet) (July 1992)